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Applicant: MITSUI TOATSU CHEMICALS, INCORPORATED
 2-5, 3-chome, Kasumigaseki Chiyoda-ku Tokyo(JP)

2 Inventor: Tanaka, Eishi 307 Detomu-1 Kyodaimae 45 Tanakaoicho Sakyo-ku Kyoto-shi Kyoto-fu(JP) Inventor: Nishizawa, Tsutomu

460-23-4-302 Kamigocho Sakae-ku Yokohama-shi Kanagawa-ken(JP) Inventor: Yamada, Yasuyuki

2-37 Mitsui Toatsu Apaato 2882 Iljimacho, Sakae-ku

Yokohama-shi Kanagawa-ken(JP)

Inventor: itoh, Hisato

2-18 Mitsui Toatsu Apaato 2882 Iljimacho, Sakae-ku

Yokohama-shi Kanagawa-ken(JP)
Inventor: Yamaguchi, Akihiro

1-13-24 Zaimokuza

Kamakura-shi Kanagawa-ken(JP)

Inventor: Nakatsuka, Masakatsu

1-11 Mitsui Toatsu Apaato 2882 lijimacho,

Sakae-ku

Yokohama-shi Kanagawa-ken(JP)

(2) Representative: Griffin, Kenneth David et al Saunders & Dolleymore 9, Rickmansworth Road

Watford Hertfordshire WD1 7HE(GB)

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Tetraphenylthiophene derivative and electrophotographic photoreceptor containing same.

There are here provided a novel tetraphenylthiophene derivative represented by the general formula

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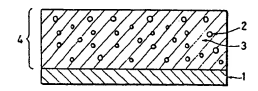
$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix} N \end{pmatrix}_{n} - \begin{pmatrix} N \\ R_2 \end{pmatrix}_{m}$$

$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix} N \end{pmatrix}_{\ell} - \begin{pmatrix} N \\ R_2 \end{pmatrix}_{m}$$

wherein each of R_1 and R_2 is an alkyl group, an aralkyl group or an aryl group which may be substituted, and R_2 and R_2 may be linked with each other to form a ring together with a nitrogen atom; and each of ℓ , madd n is an integer of 0 or 1 and is selected to meet the condition formula $\ell \ge m \ge n$;

and an electrophotographic photoreceptor which is composed of an electrically conductive base and a photosensitive layer thereon containing the aforesaid tetraphenylthiophene derivative as a charge-transporting material, and a charge-generating material, preferably a specific azo compound.

F 1 G.1



Tetraphenylthiophene Derivative and Electrophotographic Photoreceptor Containing Same

BACKGROUND OF THE INVENTION

(1) Field of the Invention

The present invention relates to a novel tetraphenylthiophene derivative and an electrophotographic photoreceptor containing the derivative. More specifically, it relates to an electrophotographic photoreceptor in which a novel tetraphenylthiophene derivative is contained as a charge-transporting material in a photosensitive layer on an electrically conductive base.

(2) Description of the Prior Art

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Heretofore, as photosensitive materials for electrophotographic photoreceptors, there have been widely used inorganic photosensitive materials such as selenium, cadmium sulfide and zinc oxide. However, the photoreceptors using such photosensitive materials do not sufficiently satisfy requirements for the electrophotographic photoreceptors such as sensitivity, light stability, moisture-proofness and durability. For example, the photoreceptors using selenium materials have excellent sensitivity but are liable to crystallize owing to heat or contaminants, so that characteristics of the photoreceptors deteriorate. In addition, this type of photoreceptors have many drawbacks. That is, manufacturing cost is high because of the utilization of vacuum evaporation, and it is difficult to mold them into the form of a belt because of less flexibility. The photoreceptors using the cadmium sulfide material are poor in moisture-proofness and durability. Furthermore, the photoreceptors in which zinc oxide is used have unsatisfactory durability.

Now, in order to eliminate these disadvantages of the photoreceptors using the inorganic photosensitive materials, various photoreceptors based on organic photosensive materials have been investigated.

Of the photoreceptors developed to remove the above-mentioned drawbacks, function-separated photoreceptors have been recently noticed in which a charge-generating function and a charge-transporting function are separately allocated to different materials. In the function-separated photoreceptors, materials having various desirable functions can be selected from a wide range of materials and combined with one another, which permits preparing the photoreceptors having high sensitivity and sufficient durability.

Requirements for the charge-transporting material contained in the electrophotographic photoreceptor are as follows:

- (1) Possessing sufficiently high ability to receive electric charges generated from the charge-generating material.
 - (2) Being capable of promptly transporting the received charges.
- (3) Being capable of successfully transporting the charges even in low electric field so that no charges remains.

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In addition, the photoreceptor should be stable to light and heat during the repeating operation of electrification, exposure, development and transfer, and should have durability enough to obtain faithfully reproduced copy images.

As the charge-transporting materials, various compounds have been reported. For example, poly(N-vinylcarbazole) is known as a light-conductive material for long time, and the photoreceptors in which this compound is used as the charge-transporting material have been put into practice. However, this type of photoreceptors are poor in flexibility and are brittle, with the result that cracks easily take place. This fact means that their durability is so bad as not to withstand the repeating use. Furthermore, when the flexibility is improved by the use of a binder, electrophotographic properties deteriorate inconveniently.

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A low-molecular weight compound has no coating properties, and thus when a photosensitive layer is formed, such a compound is mixed generally with a binder in an arbitrary ratio. Now, many charge-transporting materials have been suggested which comprise low-molecular weight compounds. For example, hydrazone compounds have high sensitivity as the charge-transporting material (see Japanese Patent Publication Laid-open Nos. 55-46761, 55-52064, 57-58156 and 57-58157). However, these compounds tend to decompose by ozone generated during corona discharge and are unstable to light and heat. In addition,

when this kind of charge-transporting material is used, obtained images are poor in contrast or have much fog owing to the degradation of charge retension ability by the repeating use or owing to the accumulation of residual potential, though initial performance is good.

Other many charge-transforming materials have been suggested, but those which sufficiently satisfy the practical requirements for use in the electrophotographic photoreceptor are not present yet, and the development of more excellent photoreceptors is desired.

SUMMARY OF THE INVENTION

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An object of the present invention is to provide a novel compound which is useful as a charge-transporting material.

Another object of the present invention is to provide an electrophotographic photoreceptor having sufficient sensitivity and excellent durability.

These objects of the present invention can be achieved by the following tetraphenylthiophene derivative and electrophotographic photoreceptor:

The tetraphenylthiophene derivative is represented by the general formula (I)

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$${\binom{R_1}{R_2}N)_n} - {\binom{N}{R_2}}_m$$

$${\binom{R_1}{R_2}N)_{\ell}} - {\binom{N}{R_2}}_{R_2}$$

$${\binom{R_1}{R_2}N)_{\ell}} - {\binom{N}{R_2}}_{R_2}$$

$${\binom{R_1}{R_2}N)_{\ell}} - {\binom{N}{R_2}}_{R_2}$$

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wherein each of R_1 and R_2 is an alkyl group, an aralkyl group or an aryl group which may be substituted, and R_1 and R_2 may be linked with each other to form a ring together with a nitrogen atom; and each of n, m and t is an integer of 0 or 1 and is selected to meet the condition formula $t \ge m \ge n$.

The electrophotographic photoreceptor comprises an electrically conductive base and a photosensitive layer thereon containing the tetraphenylthiophene derivative represented by the general formula (I) as a charge-transporting material.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1 and 2 are sectional views illustrating constitutional examples of electrophotographic photoreceptors.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

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In the above-mentioned general formula (I), examples of R_1 and R_2 include alkyl groups, which have 1 to 8 carbon atoms and which may be branched, such as a methyl group, an ethyl group, an isopropyl group, a butyl group and an octyl group; aralkyl groups, which have 7 to 10 carbon atoms in all and 1 to 4 carbon atoms on an alkyl moiety, such as a benzyl group and a phenethyl group; and aryl groups such as a phenyl group which may be substituted and a naphthyl group which may be substituted.

In case that R₁ or R₂ is the aryl group, examples of its substituent include halogen atoms; alkyl groups having 1 to 8 carbon atoms such as a methyl group, an ethyl group, a butyl group and an octyl group; alkoxy groups having 1 to 5 carbon atoms such as a methoxy group, an ethoxy group and a butoxy group;

carboxylate groups having 1 to 6 carbon atoms such as a carbomethoxy group and a carboethoxy group; aralkyl groups having 7 to 10 carbon atoms such as a benzyl group and a phenethyl group; and aryl groups such as a phenyl group and a naphthyl group.

Examples of the group in which R_1 and R_2 are linked with the nitrogen atom to form the ring include a pyrrolidino group and a piperidino group.

Preferably, R_1 and R_2 both are the aryl groups, since they give compounds which have high sensitivity and are excellent in solubility in organic solvents.

The compound represented by the above-mentioned general formula (I) can be synthesized by, for example, the following procedure:

(1) A process of reacting an amino compound represented by the general formula (V)

$$(H_2N)_n - (NH_2)_m$$

$$(H_2N)_{\ell} - (NH_2)_m$$

wherein each of £, m and n is an integer of 0 or 1 and is selected to meet the condition formula £≥m≥n, with an alkyl paratoluenesulfonate, dialkyl sulfate, an alcohol, an alkyl halide, aryl halide or the like.

(2) Anotehr method of reacting a compound represented by the general formula (VI)

$$(x)_{n} - (x)_{m}$$

$$(x)_{\ell} - (x)_{m}$$

wherein each of 1, m and n is an integer of 0 or 1 and is selected to meet the condition formula 1≥m≥n, and X is a halogen atom, preferably a iodine atom, with an amine compound represented by the general formula (VII)

$$R_1$$
 NH (VII)

wherein R₁ and R₂ are as defined above in the forlumal (I).

Typical examples of compounds which can be used in the present invention are recited in Table 1, but these examples are not restrictive.

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Table 1 (I)

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Compound No.	l	m	п	R ₁	R ₂
1	0	0	0	- (_)	-() _{CH3}
2	0	0	0	- (-()CH3
3	0	0	0	-	-
4	. 0	0	0	-()-och3	-C_>-OCH3
5	Ó	0	0	CH3-CH3	сн3-осн3
6	0	0	0	-	
7	1	0	0	-	
8	1	0	0	-	-
9	1	0	0	- <u>-</u> -c1	-<
10	1	0	0	-()-CH ₃	-{_}-Сн3

50

Table 1 (II)

Compound	۷ .	m	n	R ₁	R ₂
11	1	0	0	-{}-осн ₃	-()-OCH3
12	1	0	0	-COOC ₂ H ₅	-COOC ₂ H
13	1	0	0	-CH ₂ -	-CH ₂ -
14	1	0	0	- <u>_</u>	CH3OCH3
15	1	0	0	-CH3	-\(_\)_CH3
16	1	0	0	-<	-
17	1	0	0	-	
18	1	0	0		
19	1	0	0	-CH3	CH3OCH3
20	1	1	0	-CH3	-CH3
21	1	1	0	-	-
22	1	1	0		cl

Table 1 (III)

5	Compound No.	P	m	n	R ₁	R ₂
10	23	1	1	0	-()-cooc ₂ H ₅	-()-cooc ₂ H ₅
	24	1	1	0	-	CH3OCH3
15	25	1	1	0	-	-
	26	1	1	1	-()CH3	-CH3
20	27	1	1	1	-	-
25	28	1	1	1	-	CH3-OCH3
	29	1	1	1	-()-cooc ₂ H ₅	-COOC ₂ H ₅
30	30	1	1	1	-	-
	31	1	1	1		

The compound of the present invention which is a charge-transporting material is combined with a charge-generating material to constitute an electrophotographic photoreceptor.

As the charge-generating material, any material can be used so long as it has an ability to generate an electric charge. Typical examples of the charge-generating material include inorganic materials such as selenium, selenium alloys, amorphous silicon and cadmium sulfide; and organic dyes and pigments of phthalocyanine, perylene, perynone, indigo, anthraquinone, cyanine, azo compounds, and the like. Above all, charge-generating materials of the phthalocyanine and azo compounds are suitable for the use in combination with the charge-transporting material of the present invention.

Furthermore, of the azo compound charge-generating materials which can provide the electrophotographic photoreceptor having excellent characteristics in combination with the charge-transporting material of the present invention, the compounds represented by the following formulae (II), (III) and (IV) are particularly preferable:

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$$A-N=N-$$

$$O_{2}$$

$$-N=N-A$$
(II)

wherein A is a coupler residuum,

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$$A-N=N-$$

$$S$$

$$O_2$$

$$-N=N-A$$
(III)

wherein A is a coupler residuum,

$$A-N=N-$$

$$N=N-A$$

$$(IV)$$

$$A-N=N-A$$

wherein A is a coupler residuum.

The reason why it is particularly desirable to use each of these compounds in combination with the charge-transporting material would be based on various factors, but the presumption of these factors is extremely difficult on a present technical level in this field. It should be here noted that the combination of the charge-transporting material of the present invention and the azo compound charge-generating material having the aforesaid structural formula (II), (III) or (IV) is also on the basis of surprising discovery.

Examples of the coupler residuum represented by the group A in the general formulae (II), (III) and (IV) include the following residua (a) to (d):

(a) Coupler residua having the general formula (VIII)

wherein X is a hydrocarbon ring which may be substituted, or a heterocyclic ring which may be substituted; and Y is $-CON(R_4)-R_3$ or $-CONH^*N=C(R_5)-R_5$ wherein R_3 is a hydrocarbon ring group which may be substituted, or a heterocyclic ring group which may be substituted, R_4 is hydrocarbon ring group which may be substituted, or a phenyl group which may be substituted, a heterocyclic ring group which may be substituted, or a styryl group which may be substituted, R_5 is hydrogen, an alkyl group which may be substituted, or a phenyl group which may be substituted, and R_5 is hydrogen, an alkyl group which may be substituted, or a phenyl group which may be substituted, and R_5 and R_6 may form a ring together with carbon atoms which link therewith.

Typical examples of X in the general formula (VIII) include hydrocarbon rings such as a naphthalene ring and an anthracene ring, and heterocyclic rings such as an indole ring, a carbazole ring, a benzocarbazole ring and a dibenzofuran ring, which are each condensed with the benzene ring having a hydroxyl group and Y.

In case that X has a substituent, examples of this substituent include halogen atoms such as a chlorine atom and a bromine atom as well as a hydroxyl group.

Examples of the cyclic group of R_3 or R_5 include hydrocarbon ring groups such as a phenyl group, a naphtyl group, an anthryl group and a pyrenyl group; and heterocyclic ring groups such as a pyridyl group, a thienyl group, a furyl group, an indolyl group, a benzofuranyl group, a carbazolyl group and a

dibenzofuranyl group. Moreover, another example of the ring which can be formed by the linkage of R₅ with R₆ is a fluorene ring.

In case that R_3 or R_5 has a substituent, examples of this substituent include alkyl groups such as a methyl group, an ethyl group, a propyl group and a butyl group, alkoxy groups such as a methoxy group, an ethoxy group, a propoxy group and a butoxy group, halogen atoms such as a chlorine atom and a bromine atom, a halomethyl group such as trifluoromethyl group, dialkylamino groups such as a dimethylamino group and a diethylamino group, a nitro group, a cyano group, a carboxyl group and esters thereof.

. When R_4 or R_6 is the phenyl group, an example of its substituent is a halogen atom such as a chlorine atom or a bromine atom.

(b) Coupler residua having general formulae (IX) and (X)

wherein R₇ is a hydrocarbon group which may be substituted.

Typical examples of R₇ include alkyl groups such as a methyl group, an ethyl group, a propyl group, a butyl group and an octyl group, and alkoxyalkyl groups such as a methoxyethyl group and an ethoxyethyl group.

(c) Coupler residues having general formula (XI)

wherein R₈ is an alkyl group, a cabamoyl group, a carboxy group or its ester, and R₉ is a hydrocarbon ring group which may be substituted.

Typical examples of R_9 include hydrocarbon ring groups such as a phenyl group and a naphthyl group. In case that the group of R_9 has a substituent, examples of this substituent include alkyl groups such as a methyl group, an ethyl group, a propyl group and a buty group, alkoxy groups such as a methoxy group, an ethoxy group, a propoxy group and a butoxy group, dialkylamino groups such as a dimethylamino group and a diethylamino group, halogen atoms such as a chlorine atom and a bromine atom, a nitro group and a cyano group.

(d) Coupler residues having general formula (XII) and (XIII)

wherein Z is a divalent hydrocarbon ring group which may be substituted, or a divalent heterocyclic ring group which may be substituted.

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Typical examples of Z include a divalent monocyclic aromatic hydrocarbon group such as an ophenylene group, condensed polycyclic aromatic hydrocarbon groups such as an o-naphthylene group, a peri-naphthylene group, a 1,2-anthraquinonylene group and a 9,10-phenanthrylene group, and divalent heterocyclic ring groups such as a 3,4-pyrazoldiyl group, a 2,3-pyridinediyl group, a 4,5-pyrimidinediyl group, a 6,7-imidazolediyl group, a 5,6-benzimidazolediyl group and a 6,7-quinolinediyl group. When the ring group of Z has a substituent, examples of this substituent include alkyl groups such as a methyl group, an ethyl group, a propyl group and a butyl group, alkoxy groups such as a methoxy group, an ethoxy group, a propoxy group and a butoxy group, dialkylamino groups such as a dimethylamino group and a diethylamino group, halogen atoms such as a chlorine atom and a bromine atom, a nitro group and a cyano group.

Of the coupler residua above recited, those which are represented by the general formula (VIII) are most preferable and they have high photosensitve properties and their intermediate raw materials are so easily available that they can be manufactured at low cost.

Typical examples of the coupler residues are as follows:

HO CONH

TO CH 3

HO CONH—

A-2

25

A-3 HO CONH—CH₃

35 CH₃

HO CONH CH₃

50

12

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10

A-5

A-6

СОИН но

5 A-9

HO CONH CT

٠5

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20 A-10 HO CONH C_4H_9 (n)

25

30

A-11

HO CONH

35

10

45 A-12

HO CONH — Br

50

A-13

Br

A-17

10

15 HO CONH — CN

25
CN
HO CONH —

35

5 A-20

50

55

16

A-21

CONH . но

CONH 'HO 5 A-25

15

10

25

35

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20 A-26.

3**0** A-27

CH3 CONH 45 A-28

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A-29

15

²⁰ A-30

25

30 A-31

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40 A-32

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HO CONH -C1

Para A-34

OCH 3

HO CONH—
Br

30 EO CONH — COOH

35 40

HO CONH \longrightarrow OC₃H₇(n)

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20

5**0**

15

A-38

25

30 A-39 HO CONH OCH 3

35

40

45 A-40

HO CONH

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50

A-44

35

		CH3
5	A-45	HO CONH
10		
15		HO CONH CH3
20	A-46	H _N — O
25		
30	A-47	HO CONH—CH ³
35		HN CH
40		HO CONH—CH ₃
45	A-48	H
50		(U)

5**5**

5	A-53	HO CONH—CT
15		HO CONH $C_3H_7(n)$
20	A-54	
25		Br
30	A−55	HO CONH
35		H _M
40	·	HO CONH Br
45	A-56	H ^N
50		

Br CONH НО 5 A-57 :0 ٠5 CONH НО 20 A-58 25 CONH НО 30 A-59 35 CF₃ **≟**0 CONH НО A-60 **4**5

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A-61

A- 62

30 A-63

A-64

5	A-65	HO CONH — NO 2
15		HO CONH
20	A-66	H O
25		OCH 3
30	A-67	$\begin{array}{c} HO \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ $
35		CH ³
40		HO CONH CH3
45	A-68	, N
		•

	٠	OCH ₃
5	A-69	HO CONH —OCH 3
10		N O
15		OC_2H_5 HO CONH OC_2H_5
20	A-70	H N O
25		
30	A-71	HO CONH —COOH
35	R-11	CH3
40		HO CONH C1
45	A-72	H _N —O
50		

-N(CH₃)₂ CONH НО A-73 5 10 15 CONH -A-74 H 20 25 CONH . НО 30 A-75 H 35 CONH-40 A-76 45

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.5	A-77	HO CONH OCH 3
15		OCH ₃
20	A-78	HO CONH Br
25		
- 30	A-79	HO $CONH$ $OC_3H_7(n)$
35		
40		HO CONH \longrightarrow OC ₄ H ₉ (n)
45	. A-80	H _N
50		

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5 A-89

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20 A-90

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HO CONHN=CH CT

A-91

35

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45

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HO CONHN

50 .

CONHN=CH HO. 5 A-97 10 15 СОИНИ=СН-НО 20 A-98 25 30 HO A-99 Н 35 40 CONHN=CH НО **4**5 A-100 50

CONHN=CH НО 5 A-101 10 CONH 15 A-102 20 25 A-103 35 A-104

50

45

A-105

CON
CH
CH
3

HO CON CH₃

CH₃ OCH₃

CH₃ OCH₃

25 HO CON — CON —

HO CONHN≦CH → CONHNA€CH → CONHNA€CH

50

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38

45

соини=сн НО A-109 10 15 A-110 20 25 CONH 30 A-111 35 40 HO A-112 45 HN 50

5 A-117

10

15

A-118

25

30 A-119

A-120

45

35

50

A-121

A-122

0

A-123

A-124

5

A-125

10

15

20 A-126

HO N CH3

25

30 A-127

HO N CH³

35

40

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A-128

HO N

50

5 A-129 HO N CH₃

20 A-130

30 A-131 HO N N COOCH 3

35

40 COOH
HO N HO N

50

55

44

5 A-133 .

HO N NO 2

HO N N

A-134

20

10

15

25

A-135

35

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40 A-136

HO N CH3

50

45

5 A-137

A-138

A-139

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A-141

A-142

HO A-143

A-144

5 A-145 NO

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10

25

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A-149

10

15

A-150 20

25

30 A-151

35

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A-152

CONH НО

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The azo compounds represented by the general formula (II), (III) and (IV) can be manufactured by, for example, a method disclosed in EP 0270685.

The compound of the present invention has no ability to provide coating films by itself, and therefore it is used together with a binder in order to form a photosensitive layer. As the binder, an insulating polymer can be employed. Examples of the insulating polymer include polystyrene, polyacrylamide, polyvinyl chloride, polyester resins, polycarbonate resins, epoxy resins, phenoxy resins and polyamide resins. In particular, the polyester resins and polycarbonate resins can be suitably used. In addition, poly-N-vinylcarbazole which has charge-transporting ability by itself can be also employed.

The photoreceptor may be in the form of a constitution in which a photosensitive layer 4 containing a charge-generating material 2 and a charge-transporting material 3 is superposed upon an electrically conductive base 1, as shown in Fig. 1; a constitution in which a photosensitive layer 4 is superposed upon the electrically conductive base 1, the photosensitive layer 4 being composed of a charge-generating layer 6 containing the charge-generating material 2 and a charge-transporting layer 5 containing the charge-transporting material 3 thereon, as shown in Fig. 2; and a constitution in which the charge-generating layer 6 and the charge-transporting layer 5 in Fig. 2 are inversely disposed. The photoreceptor having any one of the above-mentioned constitutions is effective in the present invention, but the laminate type photoreceptor shown in Fig. 2 is preferble in that excellent electrophotographic characteristics can be obtained.

The constitution of the photoreceptor will be further described in detail in reference to Fig. 2.

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Examples of usable raw materials for the electrically conductive base 1 include metallic plates of aluminum, copper and zinc; those which have been prepared by depositing conductive materials such as aluminum and SnO₂ on plastic sheets or films of polyester and the like; and papers or resins which have been rendered conductive.

The charge-generating layer 6 can be formed from the above-mentioned charge-generating material 2 in accordance with a method of vacuum-evaporating the charge-generating material 2 on the electrically conductive base 1, a method of applying and drying a solution containing the charge-generating material 2, a method of applying and drying a dispersion in which fine particles of the charge-generating material 2 are dispersed, or another method.

The thickness of the charge-generating layer 6 is in the range of 0.01 to 5 μ m, preferably 0.05 to 2 μ m. When this thickness is less than 0.01 μ m, the generation of the charge is insufficient, and when it is more than 5 μ m, residual potential is high, which is practically unpreferable.

The charge-transporting layer 5 may be formed by dissolving the compound of the present invention and the above-mentioned binder in a suitable solvent, and then applying and drying the mixture. In the charge-transporting layer 5, the charge-transporting material 3 is contained in an amount of 10 to 95% by weight, preferably 30 to 90% by weight. When the charge-transporting material 3 is less than 10% by weight, the charge is scarcely transported, and when it is more than 95% by weight, mechanical strength of the photoreceptor is poor, which is practically unpreferable.

The thickness of the charge-transporting layer 5 is preferably in the range of 3 to 50 μ m, more preferably 5 to 30 μ m. When this thickness is less than 3 μ m, electrification quantity is insufficient, and when it is more than 50 μ m, residual potential is high, which is practically unpreferable.

Between the photosensitive layer and the electrically conductive base, an intermediate layer may be interposed. Suitable examples of raw materials for the intermediate layer include polyamide, nitrocellulose, casein and polyvinyl alcohol, and the thickness of the intermediate layer is preferably 1 µm or less.

As described above, the electrophotographic photoreceptor of the present invention may be composed of the compound represented by the general formula (I), the electrically conductive base, the charge-generating material and the binder, but other constitutional elements of the photoreceptor may be additionally used, so long as they have functions as such elements.

In the electrophotographic photoreceptor of the present invention, a novel compound represented by the general formula (I) is used as the charge-transporting material, and therefore this photoreceptor has high sensitivity, sufficient durability to repeating use, and other excellent advantages.

Now, the present invention will be described in detail by way of examples, but the scope of the present invention should not be limited to these examples.

Preparation Example 1

(Synthesis of Exemplary Compound No. 7)

In 50 ml of sulfolane were dispersed 12 g of 2,5-bis(4-iodophenyl)-3,4-diphenylthiophene, 10 g of N-phenyl-2-naphthylamine, 10 g of anhydrous potassium carbonate and 8 g of electrolytic copper, and the mixture was then stirred at 190 °C for 40 hours. After standing, the mixture was added to 100 ml of water, and the resultant precipitate was collected by filtration and was then washed with water and methanol in this order, followed by drying. The thus obtained crude product was extracted with benzene, and the extract was

purified through a silica gel column chromatograph using a mixed solvent of benzene and hexane (1:1) as a developing solution. By recrystallization from a mixed solvent of benzene and acetonitrile, 15 g of light-yellow crystals (melting point = 145 to 146°C) was obtained.

It was confirmed by elemental analysis that the product was Exemplary Compound No. 7.

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Results of Elemental Analysis (%)												
	C H N S											
Calcd. Found	87.56 87.34	5.14 5.08	3.40 3.31	3.90 4.04								

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Preparation Example 2

(Synthesis of Exemplary Compound No. 3)

In 200 ml of orthodichlorobenzene (hereinafter referred to simply as ODCB) was dissolved 4.1 g of 2-(4-aminophenyl)-3,4,5-triphenylthiophene, and to the mixture were added 5.1 g of iodobenzene, 11 g of anhydrous potassium carbonate, 2.6 g of electrolytic copper and 0.6 g of 18-crown-6-ether. In a nitrogen atmosphere, the mixture was then heated under reflux with vigorous stirring for 48 hours. A solid was collected by filtration, and afterward ODCB was distilled off from the reaction solution by means of steam distillation. The thus obtained residue was purified through a silica gel column chromatograph using tenzene as a developing solution. Afterward, by recrystallization from a mixed solvent of benzene and hezane (1:1), 2.5 g of light-yellow crystals (melting point = 225.0 to 228.8 °C) was obtained.

It was confirmed by elemental analysis that the product was Exemplary Compound No. 3.

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Results of Elemental Analysis (%)											
	С	Н	N	S							
Calcd. Found	86.45 86.15	5.26 5.00	2.52 2.80	5.77 6.05							

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s Preparation Example 3

(Synthesis of Exemplary Compound No. 4)

The same procedure as in Preparation Example 2 was conducted with the exception that 4.1 g of 2-(4-aminophenyl)-3,4,5-triphenylthiophene and 8.2 g of p-iodoanisole were used, to obtain 3 g of yellow crystals (melting point = 132.8 to 134.6 °C).

It was then confirmed by elemental analysis that the product was Exemplary Compound No. 4.

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Results of Elemental Analysis (%)										
	С	Н	Ν	S						
Calcd. Found	81.92 81.90	.5.40 5.19	2.27 2.16	5.20 4.91						

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Precaration Example 4

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(Synthesis of Exemplary Compound No. 8)

The same procedure as in Preparation Example 2 was conducted with the exception that 4.2 g of 2,5-bis(4-aminophenyl)-3,4-diphenylthiophene and 12.5 g of iodobenzene were used, to obtain 4 g of light-yellow crystals [melting point = 126 to 129° C; λmax = 370 nm (CHCl₃)].

It was then confirmed by elemental analysis that the product was Exemplary Compound No. 8.

Results of Elemental Analysis (%)												
	. C H N S											
Calcd. Found	86.39 86.29	5.30 4.94	3.88 3.85	4.43 4.18								

Preparation Example 5

(Synthesis of Exemplary Compound No. 10)

The same procedure as in Preparation Example 2 was conducted with the exception that 4.2 g of 2,5-bis(4-aminophenyl)-3,4-diphenylthiophene and 13.2 g of p-iodotoluene were used, to obtain 4 g of yellow crystals (melting point = 241.4 to 243.3 °C).

It was then confirmed by elemental analysis that the product was Exemplary Compound No. 10.

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Results of Elemental Analysis (%)										
	С	Н	N	S						
Calcd. Found	86.34 86.30	5.95 5.93	3.60 3.52	4.12 4.10						

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Preparation Example 6

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(Synthesis of Exemplary Compound No. 21)

The same procedure as in Preparation Example 2 was conducted with the exception that 4.3 g of 2.3.5-tris(4-aminophenyl)-4-phenylthiophene and 18.5 g of iodobenzene were used, to obtain 4 g of yellow crystals [melting point = 238 to 242° C; λmax = 373 nm (CHCl₃)].

It was then confirmed by elemental analysis and from a molecular ion peak at 889(M+) in FD-MASS that the product was Exemplary Compound No. 21.

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Results of Elemental Analysis (%)												
	C H N S											
Calcd. Found	86.36 86.82	5.32 5.22	4.72 4.52	3.60 3.38								

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Preparation Example 7

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(Synthesis of Exemplary Compound No. 27)

The same procedure as in Preparation Example 2 was conducted with the exception that 4.3 g of tetrakis(4-aminophenyl)thiophene and 24.5 g of iodobenzene were used, to obtain 6 g of yellow crystals [melting point = 228 to 232 °C; \(\lambda \text{max} = 375 \text{ nm (CHCl3)} \)].

It was then confirmed by elemental analysis and from a molecular ion peak at 889(M+) in FD-MASS that the product was Exemplary Compound No. 27.

Results of Elemental Analysis (%)										
	С	н.	N	S						
Calcd. Found	86.33 86.62	5.34 5.08	5.30 5.04	3.03 2.89						

Example 1

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In a ball mill, 0.5 g of polyester resin (made by Toyobo Co., Ltd.; trade name "Bylon 200"), 0.5 g of a diazo dye (CG-1) represented by the structural formula

wherein A is a group having the formula

and Ph is a phenyl group,

and 50 g of tetrahydrofuran were ground and mixed, and an aluminum plate was coated with the obtained dispersion by the use of a wire bar. Afterward, the coated plate was dried at 80° C for 20 minutes to form a charge-generating layer having a thickness of about 0.5 μ m.

This charge-generating layer was then coated by the use of a wire bar with a solution in which 1 g of Exemplary Compound No. 8, 1 g of polyester resin (trade name "Bylon 200"; made by Toyobo Co., Ltd.) were dissolved in 10 g of chloroform. The layer was afterward dried at 80°C for 30 minutes to form a charge-transporting layer thereon having a thickness of about 18 μ m, thereby preparing a laminate type photoreceptor as shown in Fig. 2.

The photoreceptor was electrified by corona discharge of 6 KV from an electrostatic copy paper test device (made by Kawaguchi Denki Seisakujo Co., Ltd.; model EPA-8100), and at this time, surface potential V0 was measured. Afterward, the photoreceptor was allowed to stand in a dark place for 2 seconds, and at this time, a surface potential V_2 was measured. In succession, the specimen was irradiated with light from a halogen lamp (color temperature = 2,856° K) so that the surface illuminance of the photoreceptor might be 5 lux, and a period of time which was taken until the surface potential attained a level of 1/2 of V_2 was measured. From measurements, the value of a half-life exposure $E_{1/2}$ (lux*sec) was then calculated. Furthermore, after 10 seconds were elapsed from the start of the light irradiation, a surface potential V_{12} , i.e., a residual potential was measured. Afterward, the operation of an electrification and an exposure were repeated 1,000 times.

Examples 2 to 6

The same procedure as in Example 1 was repeated with the exception that charge-transporting materials were changed to prepare photoreceptors, and measurements were done for the same items. The

used charge-transporting materials and the measured results are set forth in Table 2.

Example 7

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The same procedure as in Example 1 was conducted with the exception that a diazo dye (CG-2) represented by the following structural formula was used as a charge-generating material to prepare a photoreceptor, and measurements were done for the same items:

$$B-N=N-$$

$$-CH=CH$$

$$-CH=CH-$$

$$-N=N-B$$

wherein B is a group having the formula:

Examples 8 to 12

The same procedure as in Example 2 was conducted with the exception that the charge-transporting material was changed to prepare photoreceptors, and measurements were then made for the same items. The used charge-transporting matrials and the measured results are set forth in Table 2.

Example 13

The same procedure as in Example 1 was conducted with the exception that a disazo dye (CG-3) represented by the structural formula

$$B-N=N-$$

$$C1$$

$$-N=N-B$$

wherein B is a group having the formula

was employed as a charge-generating material to prepare a photoreceptor, and measurements were then made for the same items.

5 Example 14

The same procedure as in Example 1 was conducted with the exception that r-phthalocyanine (CG-4) was used as a charge-generating material, to prepare a photoreceptor, and measurements were then made for the same items.

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Examples 15 to 90

Following the manner of Example 1 with the exception that other charge-generating materials and charge-transporting materials were used, photoreceptors were prepared, and measurements were then made. As charge-generating materials, there were used azo compounds represented by the general formulae (II), (III) and (IV). Combinations of the used charge-generating materials and charge-transporting materials are set forth in Table 2. In this table, the used charge-generating materials are shown by symbols of the general formulae and the above-mentioned couplers.

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Comparative Example 1

The same procedure as in Example 1 was conducted with the exception that disazo dye (CG-3) represented by the structural formula

$$B-N=N-$$

$$-N=N-B$$

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wherein B is a group having the formula

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was used as a charge-generating material and 2,5-bis(4-diethylaminophenyl)-1,3,4-oxadiazole (CT-1) was used as a charge-transporting material, to prepare a photoreceptor, and the operation of electrification exposure was repeated 1,000 times.

The measured results of Examples 1 to 90 and Comparative Example 1 are set forth in Table 2.

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5		E1/2 (lux·sec)	1.2	2.0	2.2	2.0	4.2	2.8	2.0	2.3	3.1	2.1	1.8
10		<u> </u>	00	0 [-1	77	00	77	00	00	-2	00	-2-
		V2 (V)	-1080	-980 -970	-1130	-940 -900	-1270 -1200	-1150	-965 -950	-950 -945	-960 -925	-1065 -1000	-980 -940
15	Table 2 (I)	<u> </u>	-1120 -1060	-1004 -990	-1210	-990 -940	-1310	-1230 -1220	-980 -970	-980 -970	-995 -940	-1130 -1085	-1015 -975
20	Tab	Number of Repeated Operation	10001	1000	10001	1000	1000	1000	1000	1000	1000	1000	1000
30		Charge- Trans- porting Material	Compound No. 8	Compound No. 7	Compound No. 3	Compound No. 14	Compound No. 21	Compound No. 27	Compound No. 8	Compound No. 7	Compound No. 3	Compound No. 14	Compound No. 21
35		Charge- Generat- ing Material	CG-1	Ğ	ĝo	ор	ф	о́р	CG-2	go	qo	go	go
40		Example	Example 1	Example 2	Example 3	Example 4	Example 5	Example 6	Example 7	Example 8	Example 9	Example 10	Example 11

5		E1/2 (lux·sec)	3.2 6.9	1.8	2.3	2.5	2.1	3.1	ы ы 4.0.	2.2	2.3 3.8	3.6	4.4
10		V12(V)	-20	- 2	ന ന	۰-۲	77	00	5.2	77	0 -	1.1	3.2
		V2 (V)	-1065 -1010	-795 -770	-820	-950 -940	-1100 -1060	-1090 -1060	-1170	-940 -935	-1070 -1020	-1110 -1050	-1070 -1030
15	Table 2 (II)	<u>Vo(V)</u>	-1130 -1080	-815 -810	-855 -840	-1105	-1170	-1150	-1220 -1150	086- 086-	-1150	-1170	-1150 -1095
20 25	Tabl	Number of Repeated Operation	1000	10001	1000	1 1000	1000	1000	1000	1000	1000	10001	1000
30		Charge- Trans- porting Material	Compound No. 27	Compound No. 8	Compound No. 8	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30
35		Charge- Generat- ing Material	CG-2	CG-3	CG-4	(II)- (A-1)	ф	qo	do	(II)- (A-9)	đo	фo	go
40		Example	Example 12	Example 13	Example 14	Example 15	Example 16	Example 17	Example 18	Example 19	Example 20	Example 21	Example 22

5		E1/2 (lux·sec)	2.5	2.0	3.6	3.5	2.1	2.5	3.8	3.5	2.5	2.2	4.2
10		V12(V)	77	00	-50	1.1	0 -	0 -	ا ا ئ	-1-	00	-1	1.3
		V2(V)	-980 -980	-1120	-1150	-1200 -1100	-1010	-1140	-1145 -1090	-1015 -960	-970 -975	-1050 -1040	-1140
15	e 2 (III)	V ₀ (V)	-1010 -1000	-1190	-1220	-1270	-1140	-1210	-1220	-1070 -1020	-980 -985	-1110	-1220 -1120
20 25	Table 2	Number of Repeated Operation	1000	10001	1000	1000	1000	1000	1000	1000	1000	1000	10001
30		Charge- Trans- porting Material	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24
3 5		Charge- Generat- ing Material	(II)- (A-21)	ф	qo	đo	(II)- (A-49)	go	ф	đo	(II)- (A-120)	go	go
40		Example	Example 23	Example 24	Example 25	Example 26	Example 27	Example 28	Example 29	Example 30	Example 31	Example 32	Example 33

5		E1/2 (lux·sec)	3.6	2.0	2.4	3.5	3.1	2.2	2.5	4.2	8. 4. 5. 5.	2.0	2.2
10		<u>V12(V)</u>	10	00	3.5	. 0	1-2-	٠-۲	٦٥	00	E 5	0 1	0 -
		<u>v2(v)</u>	-1110 -1050	-1150	-1060 -1010	-1000 -960	-1150 -1080	-920 -910	-1185	-990 -910	-1110 -1020	-1150	-1150 -1090
15	Table 2 (IV)	70707	-1230	-1220	-1110	-1040	-1210	-1000 -970	-1250	-1040	-1160 -1090	-1230 -1200	-1220
20	Tabl	Number of Repeated Operation	10001	10001	1000	1000	1000	1000	1000	1000	1000	1000	1000
30		Charge- Trans- porting Material	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8
35		Charge- Generat- ing Material	(II)- (A-120)	(III)- (A-1)	ф	qo	do	(III)- (A-9)	go	go	ф	(III)- (A-21)	ф
40		Example	Example 34	Example 35	Example 36	Example 37	Example 38	Example 39	Example 40	Example 41	Example 42	Example 43	Example 44

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5		E1/2 (lux·sec)	3.0	3.5	2.1	2.0.	3.6	3.2	1.9	2.2	4.8 3.8	3.6	1.8
10		<u> </u>	-2	ღ <u>ც</u>	77	00	មួស	-50	00	77	2.5	10	00
		V2(V)	-1160	-920 -890	-980 -910	-1120	-1085	-1155 -1080	-940 -940	-1070	-1120	-1130 -1020	-960 -970
15	Table 2 (V)	<u> </u>	-1210 -1180	-970 -940	-1030 -990	-1180	-1120 -1080	-1240	-1040	-1140	-1180	-1210	-995 -1010
20	Tab	Number of Repeated <u>Operation</u>	1000	1000	1000	1000	1000	1000	1000	1000	1000	10001	1000
30		Charge- Trans- porting Material	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7
35		Charge- Generat- ing Material	(III)- (A-21)	qo	(III)- (A-49)	qo	đo	ф	(III)- (A-134)	ф	ор	ор	(IV)- (A-1)
40		Example	Example 45	Example 46	Example 47	Example 48	Example 49	Example 50	Example 51	Example 52	Example 53	Example 54	Example 55

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5		E1/2	2.1	3.6	3°0 8°8	0.0	3.1	2.0	3.6 4.8	2.4	2.0	3.9	3.2
10		V12(V)	00	-2	2 5 7	77	77	00	13.1	0 17	0 [-27	ო ს
		V ₂ (V)	096- 066-	-1130	-1120	-975 -970	-1150 -1130	-1165 -1150	-1065	-965 -950	-1040 -980	-1150 -1120	-1020 -920
15	Table 2 (VI)	<u>V0(V)</u>	-1065 -1020	-1210 -1220	-1190	-1020 -1000	-1230 -1220	-1180	-1090 -1060	-980 -970	-1100 -1050	-1220 -1180	-1090 -1010
20 25	Tabl	Number of Repeated <u>Operation</u>	1000	1000	1000	1000	1000	1 000	10001	10001	1000	1000	1000
30		Charge- Trans- porting Material	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30
35		Charge- Generat- ing Material	(IV)- (A-1)	ф	ф	(IV)- (A-9)	qo	ф	go	(IV)- (A-21)	đo	ф	qo
40		Example	Example 56	Example 57	Example 58	Example 59	Example 60	Example 61	Example 62	Example 63	Example 64	Example 65	Example 66

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5		E1/2 (lux·sec)	1.7	2.2	3.9	8.6 8.6	4.4	1.8	3.2	3.2	1.0	3.8 4.2	3.5.
10		V12(V)	77	0 7	00	1-13	00	00	3.2	-2	77	ლ წ	170
	_	V ₂ (V)	-1140	-1150	-1165 -1125	-955 -930	-1050	-1100 -1050	-950 -880	-990 -890	-970 -970	-1110	-1180 -1085
15	Table 2 (VII)	<u>V₀(V)</u>	-1170 -1080	-1210 -1190	-1200	-990 -950	-1170	-1150 -1095	-980 -900	-1020 -940	-1040 -1050	-1180	-1240
20 25	Tabl	Number of Repeated <u>Operation</u>	1000	1000	1000	1000	1000	1000	10001	1000	1000	1000	1000
30		Charge- Trans- porting <u>Material</u>	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24
35		Charge- Generat- ing Material	(IV)- (A-49)	go	ф	qo	(IV)- (A-53)	qo	go	ф	(IV)- (A-66)	go	go
40		Example	Example 67	Example 68	Example 69	Example 70	Example 71	Example 72	Example 73	Example 74	Example 75	Example 76	Example 77
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5		E1/2 (lux·sec)	3.7	2.1	2.6	3.3	3.5	0.8 0.8	0.0 0.9	1.1	1.1	1.2	
10		<u> </u>	77	77	127	۰۴-	77	77	00	01	77	70	77
	~	<u>V2(V)</u>	-980 -920	-1020 -1000	-1160 -1080	-1080	-970 -910	-800	-1010	-1170	-960 -970	-1010 -1000	-900 -870
15	Table 2 (VIII)	70707	-1040 -975	-1100	-1210	-1120 -1070	-1020 -950	-890	-1120	-1220	-1030 -1050	-1110	086- 086-
20 25	Table	Number of Repeated Operation	1000	1000	1000	1000	1000	1000	1000	1000	1000	1000	1000
30		Charge- Trans- porting Material	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 24	Compound No. 30	Compound No. 7	Compound No. 8	Compound No. 7	Compound No. 8	Compound No. 7	Compound No. 8
35		Charge- Generat- ing Materia <u>l</u>	(IV)- (A-66)	(IV)- (A-122)	qo	qo	ф	(IV)- (A-153)	ĝo	(IV)- (A-154)	ф	(IV)- (A-155)	op g
49		Example	Example 78	Example 79	Example 80	Example 81	Example 82	Example 83	Example 84	Example 85	Example 86	Example 87	Example 88

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5		E1/2 (lux•sec)	1.5	L 4.	5.6
10		V12(V)	. 0 1-	00	-5 -17
15		<u>V2(V)</u>	-1050 -1010	-890 -890	-950 -890
20	Table 2 (IX)	<u>Vo(V)</u>	-1170	-940 -950	-1040 -975
25	Table	Number of Repeated <u>Operation</u>	1000	1000	1000
30		Charge- Trans- porting Material	Compound No. 7	Compound No. 8	CT-1
35		Charge- Generat- ing Material	(IV)- (A-59)	qo	CG-3
40			Example 89	Example 90	Comp. Ex. 1
45		Example	Ехамр	Ехамр	Comp.

Examples 91 to 99

The photoreceptors of the present invention prepared in Examples 1, 15, 16, 35, 36, 55, 56, 83 and 84 were used in a commercially available electrophotographic copier, and copying operation was then made. In consequence, it was found that these photoreceptors could provide fog-free clear images which were faithful to an original even on the ten thousandth sheet of the copies.

Claims

1. A tetraphenylthiophene derivative represented by the general formula (I)

$$\begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} N \end{pmatrix}_{n} - \begin{pmatrix}
R_1 \\
R_2
\end{pmatrix}_{m}$$

$$\begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} N \end{pmatrix}_{\ell} - \begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} - \begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} = \begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} - \begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} = \begin{pmatrix}
R_1$$

wherein each of R_1 and R_2 is an alkyl group, an aralkyl group or an aryl group which may be substituted, and R_1 and R_2 may be linked with each other to form a ring together with a nitrogen atom; and each of ℓ , m and n is an integer of 0 or 1 and is selected to meet the condition formula $\ell \ge m \ge n$.

- 2. A tetraphenylthiophene derivative as claimed in Claim 1 wherein each of R₁ and R₂ in the formula (I) is a phenyl group which may be substituted or a naphthyl group which may be substituted.
- 3. A tetraphenylthiophene derivative as claimed in Claim 1 wherein each of R₁ and R₂ in the formula (I) is a phenyl group or a naphtyl group which may be substituted by one or more of halogen atoms, alkyl groups having 1 to 8 carbon atoms, alkoxy groups having 1 to 5 carbon atoms, carboxylate groups having 1 to 6 carbon atoms, aralkyl groups having 7 to 10 carbon atoms, phenyl groups and/or naphthyl groups.
- 4. A tetraphenylthiophene derivative as claimed in Claim 3 wherein 1, m and n in the general formula (i) are 0.
 - 5. A tetraphenylthiophene derivative as claimed in Claim 3 wherein £ is 1, and m and n are 0.
 - 6. A tetraphenylthiophene derivative as claimed in Claim 3 wherein 1 and m are 1, and n is 0.
 - 7. A tetraphenylthiophene derivative as claimed in Claim 3 wherein £, m and n are 1.
- 8. An electrophotographic photoreceptor which is composed of an electrically conductive base and a photosensitive layer thereon containing a charge-generating material and a charge-transporting material, characterized in that said charge-transporting material comprises a tetraphenylthiophene derivative represented by the general formula (I)

$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix}_{N} \begin{pmatrix} R_1 \\ R_2 \end{pmatrix}_{M} \begin{pmatrix} R_1 \\ R_2 \end{pmatrix}_{N} \begin{pmatrix} R_1 \\ R_$$

wherein each of R_1 and R_2 is an alkyl group, and an aralkyl group or an aryl group which may be substituted, R_1 and R_2 may be linked with each other to form a ring together with a nitrogen atom; and each of ℓ , m and n is an integer of 0 or 1 and is selected to meet the condition formula $\ell \ge m \ge n$.

9. An electrophotographic photoreceptor as claimed in Claim 8 wherein the photosensitive layer comprises two layers, one of which contains a charge-generating material and the other of which contains a charge-transporting material.

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- 10. An electrophotographic photoreceptor as claimed in Claim 8 wherein each of R₁ and R₂ in the formula (I) is a phenyl group or a naphtyl group which may be substituted by one or more of halogen atoms, alkyl groups having 1 to 8 carbon atoms, alkoxy groups having 1 to 5 carbon atoms, carboxylate groups having 1 to 6 carbon atoms, aralkyl groups having 7 to 10 carbon atoms, phenyl groups and/or naphthyl groups.
- 11. An electrophotographic photoreceptor as claimed in Claim 8 wherein said charge-generating material is an azo compound represented by the general formula (II), (III) or (IV)

15
$$A-N=N-$$

$$O_{2}$$

$$A-N=N-A$$

$$O_{2}$$

$$A-N=N-A$$

$$O_{2}$$

$$O_{3}$$

$$O_{4}$$

$$O_{4}$$

$$O_{5}$$

$$O_{6}$$

$$O_{7}$$

$$O_{8}$$

$$O_{9}$$

$$O_{8}$$

$$O_{9}$$

$$O_{9}$$

$$O_{1}$$

$$O_{1}$$

$$O_{1}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{3}$$

$$O_{1}$$

$$O_{1}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{1}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{3}$$

$$O_{1}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{3}$$

$$O_{4}$$

$$O_{1}$$

$$O_{1}$$

$$O_{1}$$

$$O_{2}$$

$$O_{3}$$

$$O_{4}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{1}$$

$$O_{2}$$

$$O_{3}$$

$$O_{4}$$

$$O_{5}$$

$$O_{6}$$

$$O_{7}$$

$$O_{8}$$

$$O_{9}$$

$$O_{8}$$

$$O_{8}$$

$$O_{8}$$

$$O_{8}$$

$$O_{8}$$

$$O_{8}$$

$$O_{8}$$

$$O_{9$$

wherein A in the formulae (II), (III) and (IV) is a coupler residuum.

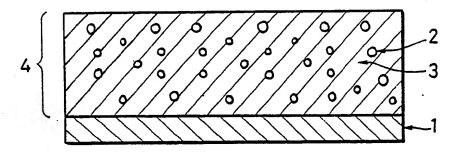
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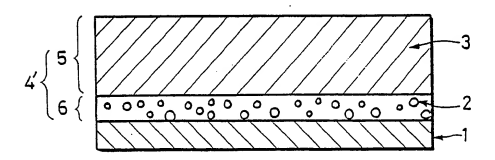
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F I G.1



F I G.2



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(71) Applicant: MITSUI TOATSU CHEMICALS. INCORPORATED 2-5, 3-chome, Kasumigaseki Chiyoda-ku Tokyo(JP)

(72) Inventor: Tanaka, Eishi 307 Detomu-1 Kyodaimae 45 Tanakaoicho Sakyo-ku Kyoto-shi Kyoto-fu(JP)

Inventor: Nishizawa, Tsutomu 460-23-4-302 Kamigocho Sakae-ku Yokohama-shi Kanagawa-ken(JP) inventor: Yamada, Yasuyuki

2-37 Mitsui Toatsu Apaato 2882 lijimacho, Sakae-ku

Yokohama-shi Kanagawa-ken(JP)

Inventor: Itoh, Hisato

2-18 Mitsui Toatsu Apaato 2882 lijimacho, Sakae-ku

Yokohama-shi Kanagawa-ken(JP) Inventor: Yamaquchi, Akihiro

1-13-24 Zaimokuza

Kamakura-shi Kanagawa-ken(JP)

Inventor: Nakatsuka, Masakatsu 1-11 Mitsui Toatsu Apaato 2882 lijimacho,

Yokohama-shi Kanagawa-ken(JP)

(4) Representative: Griffin, Kenneth David et al Saunders & Dolleymore 9, Rickmansworth Road Watford Hertfordshire WD1 7HE(GB)

- (s) Tetraphenylthiophene derivative and electrophotographic photoreceptor containing same.
- There are here provided a novel tetraphenylthiophene derivative represented by the general formula

$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix} N \end{pmatrix}_{n} - \begin{pmatrix} N \\ R_2 \end{pmatrix}$$

$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix} N \end{pmatrix}_{\ell} - \begin{pmatrix} N \\ R_2 \end{pmatrix}$$

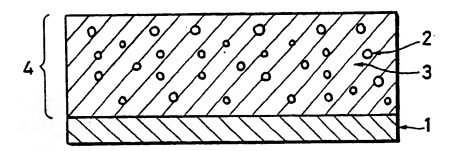
$$\begin{pmatrix} R_1 \\ R_2 \end{pmatrix} N \end{pmatrix}_{\ell} - \begin{pmatrix} N \\ R_2 \end{pmatrix}$$

wherein each of R₁ and R₂ is an alkyl group, an aralkyl group or an aryl group which may be substituted, and R1 and R2 may be linked with each other to form a ring together with a nitrogen atom; and each of 1, m and n is an integer of 0 or 1 and is selected to meet the condition formula t≥m≥n;

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and an electrophotographic photoreceptor which is composed of an electrically conductive base and a photosensitive layer thereon containing the aforesaid tetraphenylthiophene derivative as a charge-transporting material, and a charge-generating material, preferably a specific azo compound.

F I G.1





EUROPEAN SEARCH REPORT

EP 88 31 0720

]	DOCUMENTS CONSI	DERED TO BE RELEVAN	T				
Category	Citation of document with in of relevant pas	dication, where appropriate, sages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl. 4)			
Y	CHEMICAL ABSTRACTS, 5th October 1987, pa abstract no. 1245342 US; & JP-A-62 47 053 CO., LTD) 28-02-1987 * Abstract, especia	ages 585-586, z, Columbus, Ohio, 3 (FUJI PHOTO FILM	1,8,9,	C 07 D 333/20 G 03 G 5/06			
P,Y	EP-A-0 270 685 (MITCHEMICALS, INC.) * Claims *	TSUI TOATSU	1,8,9,				
P,Y	CHEMICAL ABSTRACTS, 12th December 1988, no. 211606j, Columbo JP-A-63 91 382 (ASAF 22-04-1988 * Abstract *	page 8, abstract us, Ohio, US; &	1				
				TECHNICAL FIELDS			
				SEARCHED (Int. Cl.4)			
				C 07 D 333/00 G 03 G 5/00			
	The present search report has b	een drawn up for all claims	-				
	Place of search	Date of completion of the search	01101	Examiner			
TH	E HAGUE	18-09-1990	CHO	ULY J.			
Y:pa do A:tec O:no	CATEGORY OF CITED DOCUME rticularly relevant if taken alone cument of the same category chnological background no-written disclosure termediate document	E: earlier patent de after the filing on ther D: document cited L: document cited	T: theory or principle underlying the invention E: earlier patent document, but published on, or after the filing date D: document cited in the application L: document cited for other reasons d: member of the same patent family, corresponding				

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